

		Norm	Raw		ā	%Query		:			From	To Pos	To Pos From To Pos			-
Alian Alian	Alian Alian	Align 2	`	%IDs	Align %IDs % Struct	Sea	Pairwise	Pairwise Solvation	Net	%Confidence	Pos for	į	Pos for		Alignment	Local=0
Score	Score Score	Score			Aligned	Aligned	Energy	Energy	Score	Query Query Target Target	Query	Ogen I	Target	arget	Length	Global=1
1494B800 38.21 60 (3 (align)				37.8	37.8 31.5	46.2	-17.99	-0.17	-0.957	84	30	99	20	55	37	0
149cBB00 36.91 59 (align)	59		<del>``</del>	37.8 29.4		46.2	-18.54	0.08	-0.953	83	9 08	99	28	83	37	0
ld9cA400 36.84 59 biopendium (align)	59			37.8 28.9	28.9	46.2	-17.34	1.14	-0.951	85	30	99	30	65	37	0
ingaA00 biopendium (align)	99			37.8 29.4	29.4	46.2	-8.35	09:0	-0.940	82	30	99	30	65	37	0
lotaAA00 blogendium (align)	56			31.9	15.5	59.0	-15.63	-0.64	-0.922	2. 2.	21 (6		190	234	47	0

<<Pre><<Pre>revious 1 Next>>

# Alignment

Local

Alignment Type:

Gap Open Penalty:

Segence B Range:

Seqence A Range:

```
IPAAA445 mtspnelnklpwtnpgeteicdlsdtefkISVLKNLKEIQDNTEKESRILSDKYKKQIEIIKGNQAeile
                                                                                                                                                                                                                                           10| 20| 30| 40| 50| 60
---qffreienlkeyfnggplfSEILKNWKDESDKKIIQSQIVS-FYFKLFENLKDNQVigrs
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                                      ../gtws_files/profiles/1d9gBB00.pro
                                                                                                                                               %ID
37.8
                                                                                                                                                                                                                                                                                                                                                         100|
                                                                                                                                                  Score Length Num_ID No.+ve Ovrlp
60 37 14 19 68
                                                                                                                                                                                                                                                                                                                                                         06
                                                                                                                                                                                    Lengthl Length2 Normalised-Score 111 78 38.210598
                   /usr/local/BLOSUM62
                                                       /tmp/gtw_6314.fa
                                                                                                                                                                                                                                                                                                                                                         80
                                                                                                         View Alignment: Yes
                                                                                                                             Reverse GT Alignment: No
Gap Extend Penalty:
                   Scoring Matrix:
                                    Profile A:
                                                                                        GT Alignment:
                                                       Sequence B:
                                                                        DB Alignment:
                                                                                                                                                                                                                                                                                                                                                         70
                                                                                                                                                                                                                                                                                                                                                                                                                IPAAA45 lrnadgtl---
                                                                                                                                                                                                                                                               1d9gBB00 -----
                                                                                                                                                                   SCORES:
                                                                                                                                                                                                         SCORE2:
```

Predicted sequence with translation product: INSP037 (IPAAA44548)

- 1 TGCCTAGACA CCAAAGAACA ACTATTAGCA TCAACAACAT CCAGTAAAAC ATGACTTCAC CAAACGAACT Φ ¤ Q, ល E
- 71 AAATAAGCTG CCATGGACCA ATCCTGGAGA AACAGAGATA TGTGACCTTT CAGACACAGA ATTCAAAATA 44 ರ ល ರ Ö ø u Ø b Q, ㅁ 3 Д ᅶ ㅁ
- TCTGTGTTGA AGAACCTCAA AGAAATTCAA GATAACACAG AGAAGGAATC CAGAATTCTA TCAGACAAAT Υ, ש Ø ч ល Φ × Φ ¤ ರ ש ø 7 ¤ ¥ > മ 141
- ATAAGAAACA GATTGAAATA ATTAAAGGGA ATCAAGCAGA AATTCTGGAG TTGAGAAATG CAGATGGCAC מ d ಹ ¤ ч Φ Н ø ಡ מ 디 מ ᅩ w ט ¥ Ъ, 211
- ACTITAGAAT GCATAAGAGT CTTTTTATAG CAGAATTCAT CAAGCAGAAG AAAGAAT 281

The position of primers is denoted by the reverse out boxes above.

# INSP037 (IPAAA44548) Cloned sequence with translation

- GCATCAACAA CATCCAGTAA AACATGACTT CACCAAACGA ACTAAATAAG CTGCCATGGA CCAATCCTGG l p w e l n k u മ യ
- AGAAACAGAG ATATGTGACC TTTCAGACAC AGAATTCAAA ATATCTGTGT TGAAGAACCT CAAGGAAATT **S** t e f k i յ Տ gete icd
- CAAGATAACA CAGAGAAGGA ATCCAGAATT CTATCAGACA AATATAAGAA ACAGATTGAA ATAATTAAAG 141
- GGAATCAAGC AGAAATTCTG GAGTTGAGAA ATGCAGATGG CACACTTTAG AATG e 1 r е і і 211

# Map of PCRII-TOPO-IPAAA44548

Molecule:

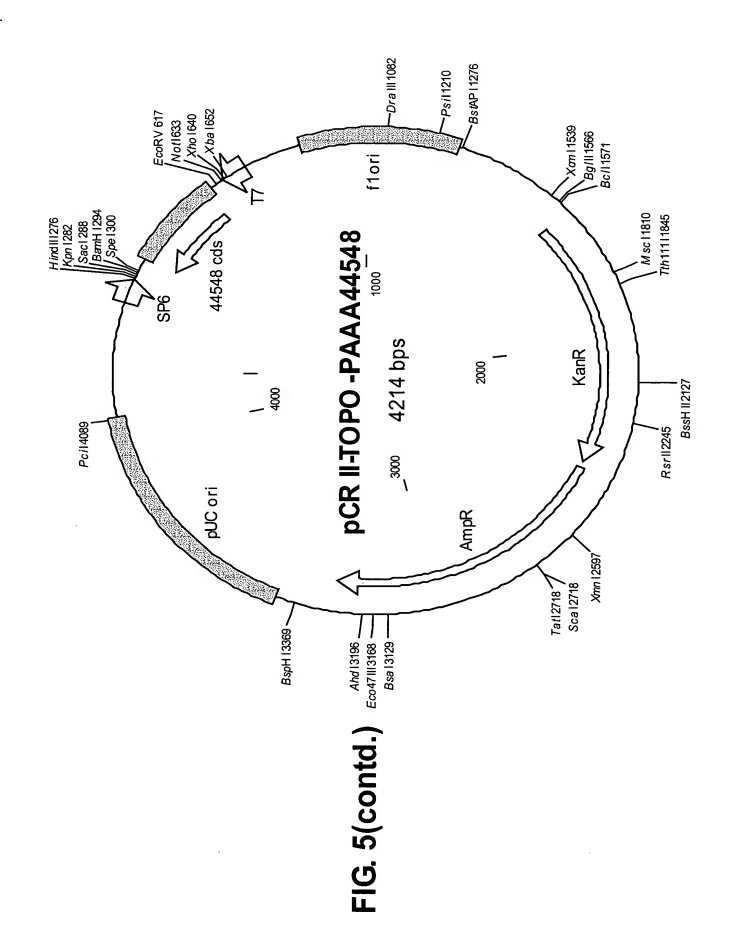
pCRII-TOPO-IPAAA44548, 4214 bps DNA Circular

File Name:

13124.cm5

Description: Plasmid ID 13124

Type	Start	End	Name	Description
MARKER	239		SP6	
REGION	337	600		IPAAA44548 cloned sequence
GENE	577	341 C	44548 cds	
MARKER	670	C	T7	
REGION	854	1268	f1 ori	
GENE	1602	2396	KanR	
GENE	2414	3274	AmpR	
REGION	3419	4092	pUC ori	
REGION GENE MARKER REGION GENE GENE	337 577 670 854 1602 2414	341 C C 1268 2396 3274	44548 cds T7 f1 ori KanR AmpR	IPAAA44548 cloned sequen



# Map of expression vector pEAK12d

Molecule:

pEAK12 d, 8760 bps DNA Circular

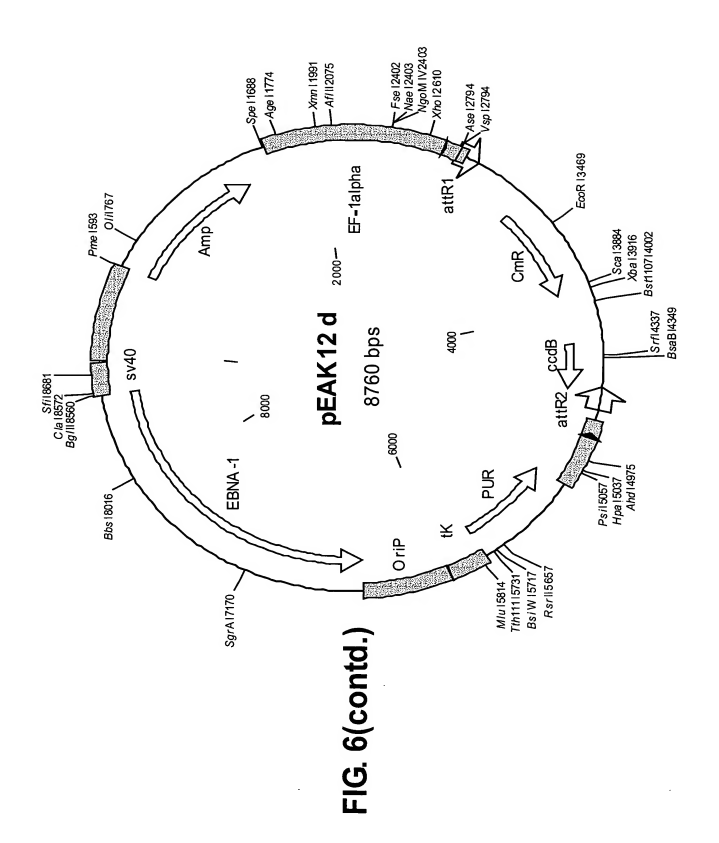
File Name:

pEAK12DEST.cm5

Description:

Mammalian cell expression vector (plasmid ID 11345)

Type	Start	End	Name	Description
REGION	2	595		pmb-ori
GENE	596	1519	Amp	•
REGION	1690	2795	EF-1alpha	
REGION	2703	2722	-	position of pEAK12F primer
REGION	2796	2845		MCS
MARKER	2855		attR1	
GENE	3256	3915	CmR	•
GENE	4257	4562	ccdB	
MARKER	4603	C	attR2	
REGION	4733	4733		MCS
REGION	4734	5162		poly A/splice
REGION	4819	4848 C		position of pEAK12R primer
GENE	5781	5163 C	PUR	PUROMYCIN
REGION	6005	5782 C	tK	tK promoter
REGION	6500	6006 C	Ori P	
GENE	8552	6500 C	EBNA-1	
REGION	8553	8752	sv40	



# Map of plasmid pDONR201

Molecule:

pDONR201, 4470 bps DNA Circular

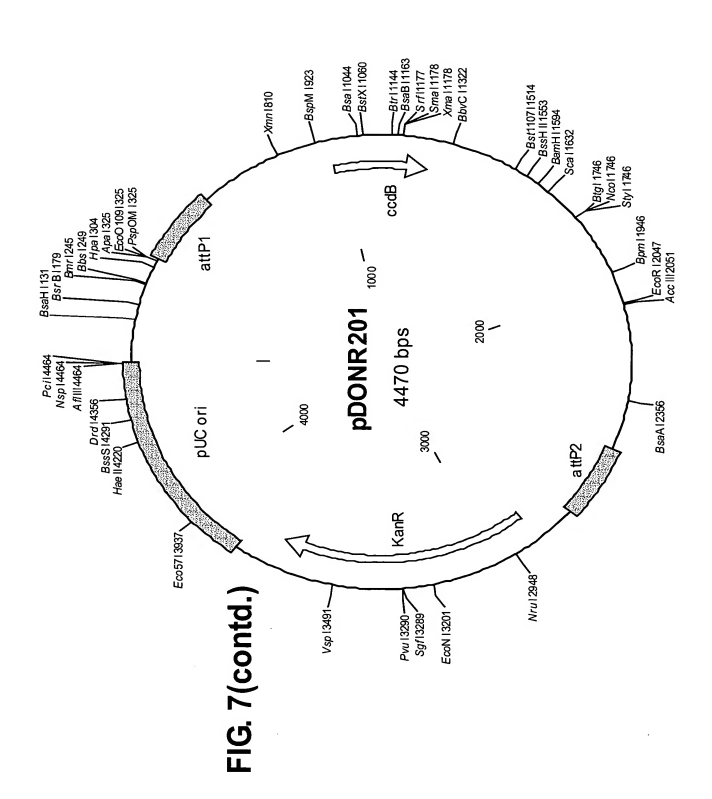
File Name:

pDONR201.cm5, dated 17 Oct 2002

Description:

Gateway entry vector (Invitrogen) - plasmid ID# 13309

Type	Start	End	Name
REGION	332	563	attP1
GENE	959	1264	ccdB
REGION	2513	2744	attP2
GENE	2868	3677	KanR
REGION	3794	4467	pUC ori



## Map of expression vector pEAK12d-IPAAA44548-6HIS

Molecule:

pEAK12d-IPAAA44548-6HIS, 7201 bps DNA Circular

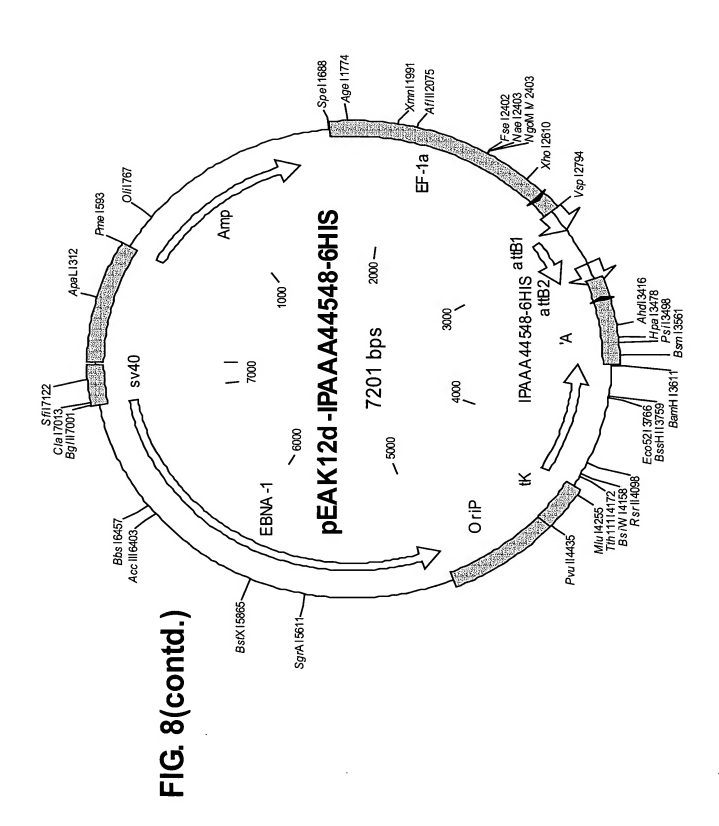
File Name:

11775.cm5

Description:

Mammalian cell Expression Construct

Туре	Start	End	Name	Description
REGION	2	595		pmb-ori
GENE	596	1519	Amp	
REGION	1690	2795	EF-1a	
REGION	2703	2722		peak12D-F primer
MARKER	2855		attB1	
GENE	2888	3139	IPAAA44548-6	HIS
MARKER	3155		attB2	
REGION	3175	3603	'A	poly A/splice
REGION	3289	3270 C		pEAK12D-R primer
GENE	4222	3604 C	•	PUROMYCIN
REGION	4446	4223 C	tK	tK promoter
REGION	4941	4447 C	Ori P	
GENE	6993	4941 C	EBNA-1	
REGION	6994	7193	sv40	



# Map of E.coli expression vector pDEST14

Molecule:

pDEST14, 6422 bps DNA Circular

File Name:

pDEST14.cm5, dated 17 Oct 2002

Description:

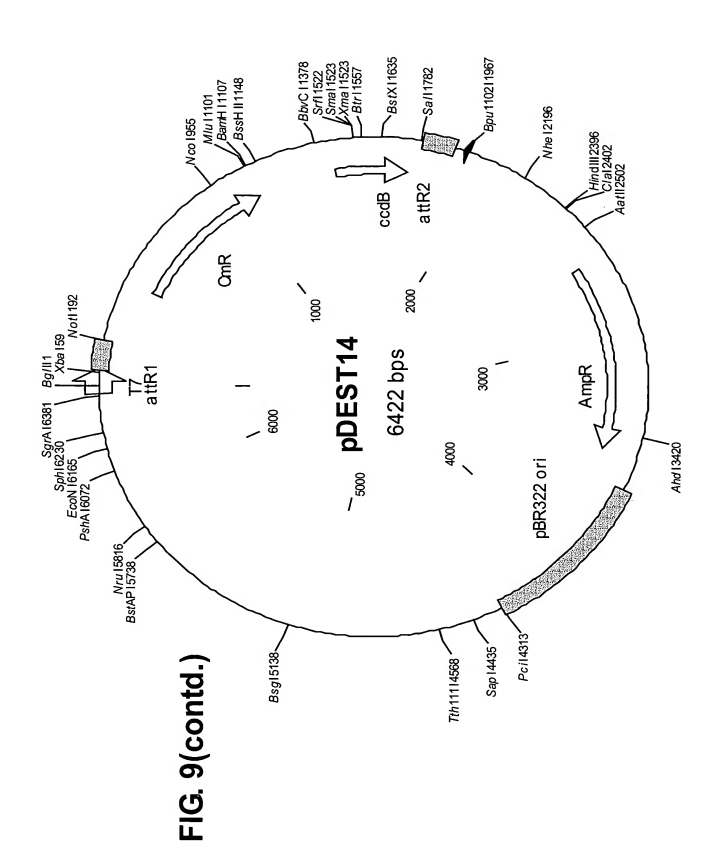
E.coli expresssion vector (Invitrogen)

Notes:

Gateway compatible, Expression under control of T7

promoter

Туре	Start	End	Name	Description
MARKER	21		T7	Promoter
REGION	67	191	attR1	
GENE	441	1100	CmR	
GENE	1442	1747	ccdB	
REGION	1788	1912	attR2	
REGION	1964	1944 C		pDEST14 R primer
GENE	2638	3498	AmpR	•
REGION	3643	4316	pBR322 ori	



# Map of plasmid pDEST14-IPAAA44548-6HIS

Molecule:

pDEST14-IPAAA44548-6HIS, 4899 bps DNA Circular

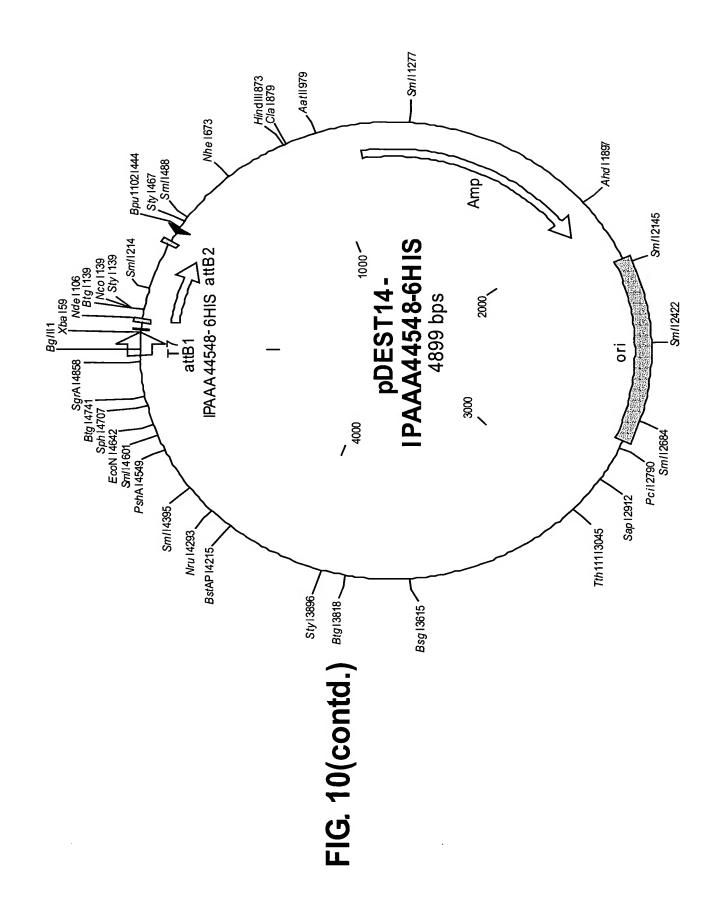
File Name:

12896.cm5

Description:

plasmid ID 12896

Туре	Start	End	Name	Description
MARKER	21		<b>T</b> 7	
REGION	72	67 C	attB1	
REGION	94	108		Shine Dalgarno Sequence
GENE	109	360	IPAAA445	48-6HIS ·
REGION	376	389	attB2	
REGION	441	421 C	1	pDEST14-R primer
GENE	1115	1975	Amp	
REGION	2124	2763	ori	pBR322 ori



#### PCRII TOPO IPAAA44548

	1	AGCGCCCAAT	ACGCAAACCG	CCTCTCCCCG	CGCGTTGGCC	GATTCATTAA	TGCAGCTGGC
	61	ACGACAGGTT	TCCCGACTGG	AAAGCGGGCA	GTGAGCGCAA	CGCAATTAAT	GTGAGTTAGC
:	121	TCACTCATTA	GGCACCCCAG	GCTTTACACT	TTATGCTTCC	GGCTCGTATG	TTGTGTGGAA
:	181	TTGTGAGCGG	ATAACAATTT	CACACAGGAA	ACAGCTATGA	CCATGATTAC	GCCAAGCTAT
2	241	TTAGGTGACA	CTATAGAATA	CTCAAGCTAT	GCATCAAGCT	TGGTACCGAG	CTCGGATCCA
:	301	CTAGTAACGG	CCGCCAGTGT	GCTGGAATTC	GCCCTTCATT	CTAAAGTGTG	CCATCTGCAT
:	361	TTCTCAACTC	CAGAATTTCT	GCTTGATTCC	CTTTAATTAT	TTCAATCTGT	TTCTTATATT
4	121	TGTCTGATAG	AATTCTGGAT	TCCTTCTCTG	TGTTATCTTG	AATTTCCTTG	AGGTTCTTCA
4	181	ACACAGATAT	TTTGAATTCT	GTGTCTGAAA	GGTCACATAT	CTCTGTTTCT	CCAGGATTGG
į	541	TCCATGGCAG	CTTATTTAGT	TCGTTTGGTG	AAGTCATGTT	TTACTGGATG	TTGTTGATGC
6	501	AAGGGCGAAT	TCTGCAGATA	TCCATCACAC	TGGCGGCCGC	TCGAGCATGC	ATCTAGAGGG
6	661	CCCAATTCGC	CCTATAGTGA	GTCGTATTAC	AATTCACTGG	CCGTCGTTTT	ACAACGTCGT
7	721	GACTGGGAAA	ACCCTGGCGT	TACCCAACTT	AATCGCCTTG	CAGCACATCC	CCCTTTCGCC
-	781	AGCTGGCGTA	ATAGCGAAGA	GGCCCGCACC	GATCGCCCTT	CCCAACAGTT	GCGCAGCCTG
8	341	AATGGCGAAT	GGGACGCGCC	CTGTAGCGGC	GCATTAAGCG	CGGCGGGTGT	GGTGGTTACG
9	901	CGCAGCGTGA	CCGCTACACT	TGCCAGCGCC	CTAGCGCCCG	CTCCTTTCGC	TTTCTTCCCT
9	961	TCCTTTCTCG	CCACGTTCGC	CGGCTTTCCC	CGTCAAGCTC	TAAATCGGGG	GCTCCCTTTA
10	21	GGGTTCCGAT	TTAGAGCTTT	ACGGCACCTC	GACCGCAAAA	AACTTGATTT	GGGTGATGGT
10	81	TCACGTAGTG	GGCCATCGCC	CTGATAGACG	GTTTTTCGCC	CTTTGACGTT	GGAGTCCACG
11	L <b>41</b>	TTCTTTAATA	GTGGACTCTT	GTTCCAAACT	GGAACAACAC	TCAACCCTAT	CGCGGTCTAT
12	201	TCTTTTGATT	TATAAGGGAT	TTTGCCGATT	TCGGCCTATT	GGTTAAAAAA	TGAGCTGATT
12	261	TAACAAATTC	AGGGCGCAAG	GGCTGCTAAA	GGAACCGGAA	CACGTAGAAA	GCCAGTCCGC
13	321	AGAAACGGTG	CTGACCCCGG	ATGAATGTCA	GCTACTGGGC	TATCTGGACA	AGGGAAAACG
13	881	CAAGCGCAAA	GAGAAAGCAG	GTAGCTTGCA	GTGGGCTTAC	ATGGCGATAG	CTAGACTGGG
14	41	CCCTTTTATC	GACAGCAAGC	GAACCGGAAT	TGCCAGCTGG	GGCGCCCTCT	GGTAAGGTTG

# FIG. 11(contd.)

1501	GGAAGCCCTG	CAAAGTAAAC	TGGATGGCTT	TCTTGCCGCC	AAGGATCTGA	TGGCGCAGGG
1561	GATCAAGATC	TGATCAAGAG	ACAGGATGAG	GATCGTTTCG	CATGATTGAA	CAAGATGGAT
1621	TGCACGCAGG	TTCTCCGGCC	GCTTGGGTGG	AGAGGCTATT	CGGCTATGAC	TGGGCACAAC
1681	AGACAATCGG	CTGCTCTGAT	GCCGCCGTGT	TCCGGCTGTC	AGCGCAGGGG	CGCCCGGTTC
1741	TTTTTGTCAA	GACCGACCTG	TCCGGTGCCC	TGAATGAACT	GCAGGACGAG	GCAGCGCGGC
1801	TATCGTGGCT	GGCCACGACG	GGCGTTCCTT	GCGCAGCTGT	GCTCGACGTT	GTCACTGAAG
1861	CGGGAAGGGA	CTGGCTGCTA	TTGGGCGAAG	TGCCGGGGCA	GGATCTCCTG	TCATCTCGCC
1921	TTGCTCCTGC	CGAGAAAGTA	TCCATCATGG	CTGATGCAAT	GCGGCGGCTG	CATACGCTTG
1981	ATCCGGCTAC	CTGCCCATTC	GACCACCAAG	CGAAACATCG	CATCGAGCGA	GCACGTACTC
2041	GGATGGAAGC	CGGTCTTGTC	GATCAGGATG	ATCTGGACGA	AGAGCATCAG	GGGCTCGCGC
2101	CAGCCGAACT	GTTCGCCAGG	CTCAAGGCGC	GCATGCCCGA	CGGCGAGGAT	CTCGTCGTGA
2161	TCCATGGCGA	TGCCTGCTTG	CCGAATATCA	TGGTGGAAAA	TGGCCGCTTT	TCTGGATTCA
2221	ACGACTGTGG	CCGGCTGGGT	GTGGCGGACC	GCTATCAGGA	CATAGCGTTG	GATACCCGTG
2281	ATATTGCTGA	AGAGCTTGGC	GGCGAATGGG	CTGACCGCTT	CCTCGTGCTT	TACGGTATCG
2341	CCGCTCCCGA	TTCGCAGCGC	ATCGCCTTCT	ATCGCCTTCT	TGACGAGTTC	TTCTGAATTG
2401	AAAAAGGAAG	AGTATGAGTA	TTCAACATTT	CCGTGTCGCC	CTTATTCCCT	TTTTTGCGGC
2461	ATTTTGCCTT	CCTGTTTTTG	CTCACCCAGA	AACGCTGGTG	AAAGTAAAAG	ATGCTGAAGA
2521	TCAGTTGGGT	GCACGAGTGG	GTTACATCGA	ACTGGATCTC	AACAGCGGTA	AGATCCTTGA
2581	GAGTTTTCGC	CCCGAAGAAC	GTTTTCCAAT	GATGAGCACT	TTTAAAGTTC	TGCTATGTGA
2641	TACACTATTA	TCCCGTATTG	ACGCCGGGCA	AGAGCAACTC	GGTCGCCGCA	TACACTATTC
2701	TCAGAATGAC	TTGGTTGAGT	ACTCACCAGT	CACAGAAAAG	CATCTTACGG	ATGGCATGAC
2761	AGTAAGAGAA	TTATGCAGTG	CTGCCATAAC	CATGAGTGAT	AACACTGCGG	CCAACTTACT
2821	TCTGACAACG	ATCGGAGGAC	CGAAGGAGCT	AACCGCTTTT	TTGCACAACA	TGGGGGATCA
2881	TGTAACTCGC	CTTGATCGTT	GGGAACCGGA	GCTGAATGAA	GCCATACCAA	ACGACGAGAG
2941	TGACACCACG	ATGCCTGTAG	CAATGCCAAC	AACGTTGCGC	AAACTATTAA	CTGGCGAACT
3001	ACTTACTCTA	GCTTCCCGGC	AACAATTAAT	AGACTGAATG	GAGGCGGATA	AAGTTGCAGG
3061	ACCACTTCTG	CGCTCGGCCC	TTCCGGCTGG	CTGGTTTATT	GCTGATAAAT	CTGGAGCCGG

3121 TGAGCGTGGG TCTCGCGGTA TCATTGCAGC ACTGGGGCCA GATGGTAAGC GCTCCCGTAT 3181 CGTAGTTATC TACACGACGG GGAGTCAGGC AACTATGGAT GAACGAAATA GACAGATCGC 3241 TGAGATAGGT GCCTCACTGA TTAAGCATTG GTAACTGTCA GACCAAGTTT ACTCATATAT 3301 ACTTTAGATT GATTTAAAAC TTCATTTTTA ATTTAAAAGG ATCTAGGTGA AGATCCTTTT 3361 TGATAATCTC ATGACCAAAA TCCCTTAACG TGAGTTTTCG TTCCACTGAG CGTCAGACCC 3421 CGTAGAAAG ATCAAAGGAT CTTCTTGAGA TCCTTTTTTT CTGCGCGTAA TCTGCTGCTT 3481 GCAAACAAAA AAACCACCGC TACCAGCGGT GGTTTGTTTG CCGGATCAAG AGCTACCAAC 3541 TCTTTTCCG AAGGTAACTG GCTTCAGCAG AGCGCAGATA CCAAATACTG TCCTTCTAGT 3601 GTAGCCGTAG TTAGGCCACC ACTTCAAGAA CTCTGTAGCA CCGCCTACAT ACCTCGCTCT 3661 GCTAATCCTG TTACCAGTGG CTGCTGCCAG TGGCGATAAG TCGTGTCTTA CCGGGTTGGA 3721 CTCAAGACGA TAGTTACCGG ATAAGGCGCA GCGGTCGGGC TGAACGGGGG GTTCGTGCAC 3781 ACAGCCCAGC TTGGAGCGAA CGACCTACAC CGAACTGAGA TACCTACAGC GTGAGCTATG 3841 AGAAAGCGCC ACGCTTCCCG AAGGGAGAAA GGCGGACAGG TATCCGGTAA GCGGCAGGGT 3901 CGGAACAGGA GAGCGCACGA GGGAGCTTCC AGGGGGAAAC GCCTGGTATC TTTATAGTCC 3961 TGTCGGGTTT CGCCACCTCT GACTTGAGCG TCGATTTTTG TGATGCTCGT CAGGGGGGCG 4021 GAGCCTATGG AAAAACGCCA GCAACGCGGC CTTTTTACGG TTCCTGGGCT TTTGCTGGCC 4081 TTTTGCTCAC ATGTTCTTTC CTGCGTTATC CCCTGATTCT GTGGATAACC GTATTACCGC 4141 CTTTGAGTGA GCTGATACCG CTCGCCGCAG CCGAACGACC GAGCGCAGCG AGTCAGTGAG 4201 CGAGGAAGCG GAAG

# FIG. 11(contd.)

#### pDEST14-IPAAA44548-6HIS

1 AGATCTCGAT CCCGCGAAAT TAATACGACT CACTATAGGG AGACCACAAC GGTTTCCCTC TAGATCACAA GTTTGTACAA 81 AAAAGCAGGC TTCGAAGGAG ATATACATAT GACTTCACCA AACGAACTAA ATAAGCTGCC ATGGACCAAT CCTGGAGAAA 161 CAGAGATATG TGACCTTTCA GACACAGAAT TCAAAATATC TGTGTTGAAG AACCTCAAGG AAATTCAAGA TAACACAGAG 241 AAGGAATCCA GAATTCTATC AGACAATAT AAGAAACAGA TTGAAATAAT TAAAGGGAAT CAAGCAGAAA TTCTGGAGTT GAGAAATGCA GATGGCACAC TTCACCATCA CCATCACCAT TGAAACCCAG CTTTCTTGTA CAAAGTGGTG ATGATCCGGC 401 TGCTAACAAA GCCCGAAAGG AAGCTGAGTT GGCTGCTGCC ACCGCTGAGC AATAACTAGC ATAACCCCTT GGGGCCTCTA 481 AACGGGTCTT GAGGGGTTTT TTGCTGAAAG GAGGAACTAT ATCCGGATAT CCACAGGACG GGTGTGGTCG CCATGATCGC GTAGTCGATA GTGGCTCCAA GTAGCGAAGC GAGCAGGACT GGGCGGCGGC CAAAGCGGTC GGACAGTGCT CCGAGAACGG 641 GTGCGCATAG AAATTGCATC AACGCATATA GCGCTAGCAG CACGCCATAG TGACTGGCGA TGCTGTCGGA ATGGACGATA 721 TCCCGCAAGA GGCCCGGCAG TACCGGCATA ACCAAGCCTA TGCCTACAGC ATCCAGGGTG ACGGTGCCGA GGATGACGAT GAGCGCATTG TTAGATTCA TACACGGTGC CTGACTGCGT TAGCAATTTA ACTGTGATAA ACTACCGCAT TAAAGCTTAT 881 CGATGATAAG CTGTCAAACA TGAGAATTCT TGAAGACGAA AGGGCCTCGT GATACGCCTA TTTTTATAGG TTAATGTCAT GATAATAATG GTTTCTTAGA CGTCAGGTGG CACTTTTCGG GGAAATGTGC GCGGAACCCC TATTTGTTTA TTTTTCTAAA TACATTCAAA TATGTATCCG CTCATGAGAC AATAACCCTG ATAAATGCTT CAATAATATT GAAAAAGGAA GAGTATGAGT 1121 ATTCAACATT TCCGTGTCGC CCTTATTCCC TTTTTTGCGG CATTTTGCCT TCCTGTTTTT GCTCACCCAG AAACGCTGGT 1201 GAAAGTAAAA GATGCTGAAG ATCAGTTGGG TGCACGAGTG GGTTACATCG AACTGGATCT CAACAGCGGT AAGATCCTTG 1281 AGAGTTTTCG CCCCGAAGAA CGTTTTCCAA TGATGAGCAC TTTTAAAGTT CTGCTATGTG GCGCGGTATT ATCCCGTGTT GACGCCGGGC AAGAGCAACT CGGTCGCCGC ATACACTATT CTCAGAATGA CTTGGTTGAG TACTCACCAG TCACAGAAAA 1441 GCATCTTACG GATGGCATGA CAGTAAGAGA ATTATGCAGT GCTGCCATAA CCATGAGTGA TAACACTGCG GCCAACTTAC 1521 TTCTGACAAC GATCGGAGGA CCGAAGGAGC TAACCGCTTT TTTGCACAAC ATGGGGGATC ATGTAACTCG CCTTGATCGT TGGGAACCGG AGCTGAATGA AGCCATACCA AACGACGAGC GTGACACCAC GATGCCTGCA GCAATGGCAA CAACGTTGCG 1681 CAAACTATTA ACTGGCGAAC TACTTACTCT AGCTTCCCGG CAACAATTAA TAGACTGGAT GGAGGCGGAT AAAGTTGCAG 1761 GACCACTTCT GCGCTCGGCC CTTCCGGCTG GCTGGTTTAT TGCTGATAAA TCTGGAGCCG GTGAGCGTGG GTCTCGCGGT 1841 ATCATTGCAG CACTGGGGCC AGATGGTAAG CCCTCCCGTA TCGTAGTTAT CTACACGACG GGGAGTCAGG CAACTATGGA TGAACGAAAT AGACAGATCG CTGAGATAGG TGCCTCACTG ATTAAGCATT GGTAACTGTC AGACCAAGTT TACTCATATA 1921 2001 TACTTTAGAT TGATTTAAAA CTTCATTTTT AATTTAAAAG GATCTAGGTG AAGATCCTTT TTGATAATCT CATGACCAAA 2081 ATCCCTTAAC GTGAGTTTTC GTTCCACTGA GCGTCAGACC CCGTAGAAAA GATCAAAGGA TCTTCTTGAG ATCCTTTTTT 2161 TCTGCGCGTA ATCTGCTGCT TGCAAACAAA AAAACCACCG CTACCAGCGG TGGTTTGTTT GCCGGATCAA GAGCTACCAA 2241 CTCTTTTTCC GAAGGTAACT GGCTTCAGCA GAGCGCAGAT ACCAAATACT GTCCTTCTAG TGTAGCCGTA GTTAGGCCAC 2321 CACTTCAAGA ACTCTGTAGC ACCGCCTACA TACCTCGCTC TGCTAATCCT GTTACCAGTG GCTGCTGCCA GTGGCGATAA

2401	GTCGTGTCTT	ACCGGGTTGG	ACTCAAGACG	ATAGTTACCG	GATAAGGCGC	AGCGGTCGGG	CTGAACGGGG	GGTTCGTGCA
2481	CACAGCCCAG	CTTGGAGCGA	ACGACCTACA	CCGAACTGAG	ATACCTACAG	CGTGAGCTAT	GAGAAAGCGC	CACGCTTCCC
2561	GAAGGGAGAA	AGGCGGACAG	GTATCCGGTA	AGCGGCAGGG	TCGGAACAGG	AGAGCGCACG	AGGGAGCTTC	CAGGGGGAAA
2641	CGCCTGGTAT	CTTTATAGTC	CTGTCGGGTT	TCGCCACCTC	TGACTTGAGC	GTCGATTTTT	GTGATGCTCG	TCAGGGGGGC
2721	GGAGCCTATG	GAAAAACGCC	AGCAACGCGG	CCTTTTTACG	GTTCCTGGCC	TTTTGCTGGC	CTTTTGCTCA	CATGTTCTTT
2801	CCTGCGTTAT	CCCCTGATTC	TGTGGATAAC	CGTATTACCG	CCTTTGAGTG	AGCTGATACC	GCTCGCCGCA	GCCGAACGAC
2881	CGAGCGCAGC	GAGTCAGTGA	GCGAGGAAGC	GGAAGAGCGC	CTGATGCGGT	ATTTTCTCCT	TACGCATCTG	TGCGGTATTT
2961	CACACCGCAT	ATATGGTGCA	CTCTCAGTAC	AATCTGCTCT	GATGCCGCAT	AGTTAAGCCA	GTATACACTC	CGCTATCGCT
3041	ACGTGACTGG	GTCATGGCTG	CGCCCCGACA	CCCGCCAACA	CCCGCTGACG	CGCCCTGACG	GGCTTGTCTG	CTCCCGGCAT
3121	CCGCTTACAG	ACAAGCTGTG	ACCGTCTCCG	GGAGCTGCAT	GTGTCAGAGG	TTTTCACCGT	CATCACCGAA	ACGCGCGAGG
3201	CAGCTGCGGT	AAAGCTCATC	AGCGTGGTCG	TGAAGCGATT	CACAGATGTC	TGCCTGTTCA	TCCGCGTCCA	GCTCGTTGAG
3281	TTTCTCCAGA	AGCGTTAATG	TCTGGCTTCT	GATAAAGCGG	GCCATGTTAA	GGGCGGTTTT	TTCCTGTTTG	GTCACTGATG
3361	CCTCCGTGTA	AGGGGGATTT	CTGTTCATGG	GGGTAATGAT	ACCGATGAAA	CGAGAGAGGA	TGCTCACGAT	ACGGGTTACT
3441	GATGATGAAC	ATGCCCGGTT	ACTGGAACGT	TGTGAGGGTA	AACAACTGGC	GGTATGGATG	CGGCGGGACC	AGAGAAAAAT
3521	CACTCAGGGT	CAATGCCAGC	GCTTCGTTAA	TACAGATGTA	GGTGTTCCAC	AGGGTAGCCA	GCAGCATCCT	GCGATGCAGA
3601	TCCGGAACAT	AATGGTGCAG	GGCGCTGACT	TCCGCGTTTC	CAGACTTTAC	GAAACACGGA	AACCGAAGAC	CATTCATGTT
3681	GTTGCTCAGG	TCGCAGACGT	TTTGCAGCAG	CAGTCGCTTC	ACGTTCGCTC	GCGTATCGGT	GATTCATTCT	GCTAACCAGT
3761	AAGGCAACCC	CGCCAGCCTA	GCCGGGTCCT	CAACGACAGG	AGCACGATCA	TGCGCACCCG	TGGCCAGGAC	CCAACGCTGC
3841	CCGAGATGCG	CCGCGTGCGG	CTGCTGGAGA	TGGCGGACGC	GATGGATATG	TTCTGCCAAG	GGTTGGTTTG	CGCATTCACA
3921	GTTCTCCGCA	AGAATTGATT	GGCTCCAATT	CTTGGAGTGG	TGAATCCGTT	AGCGAGGTGC	CGCCGGCTTC	CATTCAGGTC
4001	GAGGTGGCCC	GGCTCCATGC	ACCGCGACGC	AACGCGGGGA	GGCAGACAAG	GTATAGGGCG	GCGCCTACAA	TCCATGCCAA
4081	CCCGTTCCAT	GTGCTCGCCG	AGGCGGCATA	AATCGCCGTG	ACGATCAGCG	GTCCAGTGAT	CGAAGTTAGG	CTGGTAAGAG
4161	CCGCGAGCGA	TCCTTGAAGC	TGTCCCTGAT	GGTCGTCATC	TACCTGCCTG	GACAGCATGG	CCTGCAACGC	GGGCATCCCG
4241	ATGCCGCCGG	AAGCGAGAAG	AATCATAATG	GGGAAGGCCA	TCCAGCCTCG	CGTCGCGAAC	GCCAGCAAGA	CGTAGCCCAG
4321	CGCGTCGGCC	GCCATGCCGG	CGATAATGGC	CTGCTTCTCG	CCGAAACGTT	TGGTGGCGGG	ACCAGTGACG	AAGGCTTGAG
4401	CGAGGGCGTG	CAAGATTCCG	AATACCGCAA	GCGACAGGCC	GATCATCGTC	GCGCTCCAGC	GAAAGCGGTC	CTCGCCGAAA
4481	ATGACCCAGA	GCGCTGCCGG	CACCTGTCCT	ACGAGTTGCA	TGATAAAGAA	GACAGTCATA	AGTGCGGCGA	CGATAGTCAT
4561	GCCCGCGCC	CACCGGAAGG	AGCTGACTGG	GTTGAAGGCT	CTCAAGGGCA	TCGGTCGATC	GACGCTCTCC	CTTATGCGAC
4641	TCCTGCATTA	GGAAGCAGCC	CAGTAGTAGG	TTGAGGCCGT	TGAGCACCGC	CGCCGCAAGG	AATGGTGCAT	GCAAGGAGAT
4721	GGCGCCCAAC	AGTCCCCCGG	CCACGGGGCC	TGCCACCATA	CCCACGCCGA	AACAAGCGCT	CATGAGCCCG	AAGTGGCGAG
4801	CCCGATCTTC	CCCATCGGTG					_	GGCCACGATG
4881	CGTCCGGCGT	AGAGGATCG		FIG.	12(C	ontd.	)	

#### pEAK12D-IPAAA44548-6HIS

1 GGCGTAATCT GCTGCTTGCA AACAAAAAA CCACCGCTAC CAGCGGTGGT TTGTTTGCCG GATCAAGAGC TACCAACTCT 81 TTTTCCGAAG GTAACTGGCT TCAGCAGAGC GCAGATACCA AATACTGTCC TTCTAGTGTA GCCGTAGTTA GGCCACCACT 161 TCAAGAACTC TGTAGCACCG CCTACATACC TCGCTCTGCT GAAGCCAGTT ACCAGTGGCT GCTGCCAGTG GCGATAAGTC 241 GTGTCTTACC GGGTTGGACT CAAGAGATAG TTACCGGATA AGGCGCAGCG GTCGGGCTGA ACGGGGGGTT CGTGCACACA 321 GCCCAGCTTG GAGCGAACGA CCTACACCGA ACTGAGATAC CTACAGCGTG AGCTATGAGA AAGCGCCACG CTTCCCGAAG 401 GGAGAAAGGC GGACAGGTAT CCGGTAAGCG GCAGGGTCGG AACAGGAGAG CGCACGAGGG AGCTTCCAGG GGGAAACGCC TGGTATCTTT ATAGTCCTGT CGGGTTTCGC CACCTCTGAC TTGAGCGTCG ATTTTTGTGA TGCTCGTCAG GGGGGCGGAG 561 CCTATGGAAA AACGCCAGCA ACGCAAGCTA GAGTTTAAAC TTGACAGATG AGACAATAAC CCTGATAAAT GCTTCAATAA TATTGAAAAA GGAAAAGTAT GAGTATTCAA CATTTCCGTG TCGCCCTTAT TCCCTTTTTT GCGGCATTTT GCCTTCCTGT 641 721 TTTTGCTCAC CCAGAAACGC TGGTGAAAGT AAAAGATGCA GAAGATCACT TGGGTGCGCG AGTGGGTTAC ATCGAACTGG 801 ATCTCAACAG CGGTAAGATC CTTGAGAGTT TTCGCCCCGA AGAACGTTTC CCAATGATGA GCACTTTTAA AGTTCTGCTA TGTGGCGCGG TATTATCCCG TATTGATGCC GGGCAAGAGC AACTCGGTCG CCGCATACAC TATTCTCAGA ATGACTTGGT TGAATACTCA CCAGTCACAG AAAAGCATCT TACGGATGGC ATGACAGTAA GAGAATTATG CAGTGCTGCC ATAACCATGA 1041 GTGATAACAC TGCGGCCAAC TTACTTCTGA CAACTATCGG AGGACCGAAG GAGCTAACCG CTTTTTTGCA CAACATGGGG 1121 GATCATGTAA CTCGCCTTGA TCGTTGGGAA CCGGAGCTGA ATGAAGCCAT ACCAAACGAC GAGCGTGACA CCACGATGCC 1201 TGTAGCAATG GCAACAACGT TGCGAAAACT ATTAACTGGC GAACTACTTA CTCTAGCTTC CCGGCAACAA CTAATAGACT 1281 GGATGGAGGC GGATAAAGTT GCAGGACCAC TTCTGCGCTC GGCACTTCCG GCTGGCTGGT TTATTGCTGA TAAATCAGGA 1361 GCCGGTGAGC GTGGGTCACG CGGTATCATT GCAGCACTGG GGCCGGATGG TAAGCCCTCC CGTATCGTAG TTATCTACAC TACGGGGAGT CAGGCAACTA TGGATGAACG AAATAGACAG ATCGCTGAGA TAGGTGCCTC ACTGATTAAG CATTGGTAAG GATAAATTTC TGGTAAGGAG GACACGTATG GAAGTGGGCA AGTTGGGGGAA GCCGTATCCG TTGCTGAATC TGGCATATGT 1521 1601 GGGAGTATAA GACGCGCAGC GTCGCATCAG GCATTTTTTT CTGCGCCAAT GCAAAAAGGC CATCCGTCAG GATGGCCTTT CGGCATAACT AGTGAGGCTC CGGTGCCCGT CAGTGGGCAG AGCGCACATC GCCCACAGTC CCCGAGAAGT TGGGGGGAGG GGTCGGCAAT TGAACCGGTG CCTAGAGAAG GTGGCGCGGG GTAAACTGGG AAAGTGATGT CGTGTACTGG CTCCGCCTTT TTCCCGAGGG TGGGGGAGAA CCGTATATAA GTGCAGTAGT CGCCGTGAAC GTTCTTTTTC GCAACGGGTT TGCCGCCAGA ACACAGGTAA GTGCCGTGTG TGGTTCCCGC GGGCCTGGCC TCTTTACGGG TTATGGCCCT TGCGTGCCTT GAATTACTTC CACCTGGCTG CAGTACGTGA TTCTTGATCC CGAGCTTCGG GTTGGAAGTG GGTGGGAGAG TTCGAGGCCT TGCGCTTAAG 2161 GCCTGTCTCG CTGCTTTCGA TAAGTCTCTA GCCATTTAAA ATTTTTGATG ACCTGCTGCG ACGCTTTTTT TCTGGCAAGA

# FIG. 13(contd.)

2241 TAGTCTTGTA AATGCGGGCC AAGACGATCT GCACACTGGT ATTTCGGTTT TTGGGGCCGC GGGCGGCGAC GGGGCCCGTG 2321 CGTCCCAGCG CACATGCATG TTCGGCGAGG CGGGGCCTGC GAGCGCGGCC ACCGAGAATC GGACGGGGGT AGTCTCAAGC TGGCCGGCCT GCTCTGGTGC CTGGCCTCGC GCCGCCGTGT ATCGCCCCGC CCTGGGCGGC AAGGCTGGGA GCTCAAAATG GAGGACGCGG CGCTCGGGAG AGCGGGCGGG TGAGTCACCC ACACAAAGGA AAAGGGCCTT TCCGTCCTCA GCCGTCGCTT CATGTGACTC CACGGAGTAC CGGGCGCCGT CCAGGCACCT CGATTAGTTC TCGAGCTTTT GGAGTACGTC GTCTTTAGGT TGGGGGGAGG GGTTTTATGC GATGGAGTTT CCCCACACTG AGTGGGTGGA GACTGAAGTT AGGCCAGCTT GGCACTTGAT GTAATTCTCC TTGGAATTTG CCCTTTTTGA GTTTGGATCT TGGTTCATTC TCAAGCCTCA GACAGTGGTT CAAATTAATA CGACTCACTA TAGGGAGACT TCTTTCTCCC ATTTCAGGTG TCGTAAGCTA TCAAACAAGT TTGTACAAAA AAGCAGGCTT CGCCACCATG ACTTCACCAA ACGAACTAAA TAAGCTGCCA TGGACCAATC CTGGAGAAAC AGAGATATGT GACCTTTCAG ACACAGAATT CAAAATATCT GTGTTGAAGA ACCTCAAGGA AATTCAAGAT AACACAGAGA AGGAATCCAG AATTCTATCA GACAAATATA AGAAACAGAT TGAAATAATT AAAGGGAATC AAGCAGAAAT TCTGGAGTTG AGAAATGCAG ATGGCACACT TCACCATCAC CATCACCATT GAAACCCAGC TTTCTTGTAC AAAGTGGTTC GATGGCCGCA GGTAAGCCAG CCCAGGCCTC GCCCTCCAGC TCAAGGCGGG ACAGGTGCCC TAGAGTAGCC TGCATCCAGG GACAGGCCCC AGCCGGGTGC TGACACGTCC ACCTCCATCT CTTCCTCAGG TCTGCCCGGG TGGCATCCCT GTGACCCCTC CCCAGTGCCT CTCCTGGTCG TGGAAGGTGC TACTCCAGTG CCCACCAGCC TTGTCCTAAT AAAATTAAGT TGCATCATTT TGTTTGACTA GGTGTCCTTG TATAATATTA TGGGGTGGAG GCGGGTGGTA TGGAGCAAGG GGCCCAAGTT AACTTGTTTA TTGCAGCTTA TAATGGTTAC AAATAAAGCA 3521 ATAGCATCAC AAATTTCACA AATAAAGCAT TTTTTTCACT GCATTCTAGT TGTGGTTTGT CCAAACTCAT CAATGTATCT TATCATGTCT GGATCCGCTT CAGGCACCGG GCTTGCGGGT CATGCACCAG GTGCGCGGTC CTTCGGGCAC CTCGACGTCG GCGGTGACGG TGAAGCCGAG CCGCTCGTAG AAGGGGAGGT TGCGGGGCGC GGAGGTCTCC AGGAAGGCGG GCACCCCGGC GCGCTCGGCC GCCTCCACTC CGGGGAGCAC GACGGCGCTG CCCAGACCCT TGCCCTGGTG GTCGGGCGAG ACGCCGACGG TGGCCAGGAA CCACGCGGGC TCCTTGGGCC GGTGCGGCGC CAGGAGGCCT TCCATCTGTT GCTGCGCGGC CAGCCTGGAA CCGCTCAACT CGGCCATGCG CGGGCCGATC TCGGCGAACA CCGCCCCCGC TTCGACGCTC TCCGGCGTGG TCCAGACCGC CACCGCGGCG CCGTCGTCCG CGACCCACAC CTTGCCGATG TCGAGCCCGA CGCGCGTGAG GAAGAGTTCT TGCAGCTCGG TGACCCGCTC GATGTGGCGG TCCGGGTCGA CGGTGTGGCG CGTGGCGGGG TAGTCGGCGA ACGCGGCGGC GAGGGTGCGT ACGGCCCGGG GGACGTCGTC GCGGGTGGCG AGGCGCACCG TGGGCTTGTA CTCGGTCATG GTGGCCTGCA GAGTCGCTCT GTGTTCGAGG CCACACGCGT CACCTTAATA TGCGAAGTGG ACCTGGGACC GCGCCGCCCC GACTGCATCT GCGTGTTTTC GCCAATGACA AGACGCTGGG CGGGGTTTGT GTCATCATAG AACTAAAGAC ATGCAAATAT ATTTCTTCCG GGGACACCGC CAGCAAACGC GAGCAACGGG CCACGGGGAT GAAGCAGCTG CGCCACTCCC TGAAGATCCC CCTTATTAAC CCTAAACGGG TAGCATATGC TTCCCGGGTA GTAGTATATA CTATCCAGAC TAACCCTAAT TCAATAGCAT ATGTTACCCA ACGGGAAGCA TATGCTATCG AATTAGGGTT AGTAAAAGGG TCCTAAGGAA CAGCGATCTG GATAGCATAT GCTATCCTAA TCTATATCTG 4641 GGTAGCATAT GCTATCCTAA TCTATATCTG GGTAGCATAG GCTATCCTAA TCTATATCTG GGTAGCATAT GCTATCCTAA 4721 TCTATATCTG GGTAGTATAT GCTATCCTAA TTTATATCTG GGTAGCATAG GCTATCCTAA TCTATATCTG GGTAGCATAT

4801	GCTATCCTAA	TCTATATCTG	GGTAGTATAT	GCTATCCTAA	TCTGTATCCG	GGTAGCATAT	GCTATCCTCA	TGCATATACA
4881	GTCAGCATAT	GATACCCAGT	AGTAGAGTGG	GAGTGCTATC	CTTTGCATAT	GCCGCCACCT	CCCAAGGAGA	TCCGCATGTC
4961	TGATTGCTCA	CCAGGTAAAT	GTCGCTAATG	TTTTCCAACG	CGAGAAGGTG	TTGAGCGCGG	AGCTGAGTGA	CGTGACAACA
5041	TGGGTATGCC	CAATTGCCCC	ATGTTGGGAG	GACGAAAATG	GTGACAAGAC	AGATGGCCAG	AAATACACCA	ACAGCACGCA
5121	TGATGTCTAC	TGGGGATTTA	TTCTTTAGTG	CGGGGGAATA	CACGGCTTTT	AATACGATTG	AGGGCGTCTC	CTAACAAGTT
5201	ACATCACTCC	TGCCCTTCCT	CACCCTCATC	TCCATCACCT	CCTTCATCTC	CGTCATCTCC	GTCATCACCC	TCCGCGGCAG
5281	CCCCTTCCAC	CATAGGTGGA	AACCAGGGAG	GCAAATCTAC	TCCATCGTCA	AAGCTGCACA	CAGTCACCCT	GATATTGCAG
5361	GTAGGAGCGG	GCTTTGTCAT	AACAAGGTCC	TTAATCGCAT	CCTTCAAAAC	CTCAGCAAAT	ATATGAGTTT	GTAAAAAGAC
5441	CATGAAATAA	CAGACAATGG	ACTCCCTTAG	CGGGCCAGGT	TGTGGGCCGG	GTCCAGGGGC	CATTCCAAAG	GGGAGACGAC
5521	TCAATGGTGT	AAGACGACAT	TGTGGAATAG	CAAGGGCAGT	TCCTCGCCTT	AGGTTGTAAA	GGGAGGTCTT	ACTACCTCCA
5601	TATACGAACA	CACCGGCGAC	CCAAGTTCCT	TCGTCGGTAG	TCCTTTCTAC	GTGACTCCTA	GCCAGGAGAG	CTCTTAAACC
5681	TTCTGCAATG	TTCTCAAATT	TCGGGTTGGA	ACCTCCTTGA	CCACGATGCT	TTCCAAACCA	CCCTCCTTTT	TTGCGCCTGC
5761	CTCCATCACC	CTGACCCCGG	GGTCCAGTGC	TTGGGCCTTC	TCCTGGGTCA	TCTGCGGGGC	CCTGCTCTAT	CGCTCCCGGG
5841	GGCACGTCAG	GCTCACCATC	TGGGCCACCT	TCTTGGTGGT	ATTCAAAATA	ATCGGCTTCC	CCTACAGGGT	GGAAAAATGG
5921	CCTTCTACCT	GGAGGGGGCC	TGCGCGGTGG	AGACCCGGAT	GATGATGACT	GACTACTGGG	ACTCCTGGGC	CTCTTTTCTC
6001	CACGTCCACG	ACCTCTCCCC	CTGGCTCTTT	CACGACTTCC	CCCCTGGCT	CTTTCACGTC	CTCTACCCCG	GCGGCCTCCA
6081	CTACCTCCTC	GACCCCGGCC	TCCACTACCT	CCTCGACCCC	GGCCTCCACT	GCCTCCTCGA	CCCCGGCCTC	CGGCACCTCC
6161	TCCAGCCCCA	GCACCTCCAC	CAGCCCCAGC	TCCCCCAGCT	CCAGCCCCAC	CAGCACCAGC	CCCTCCAGCC	CCACCAGCCC
6241	CAGCCCCTCC	GGCACCTCCT	CCAGCCCCAG	CACCTCCACC	AGCCCCAGCT	CCCCCAGCTC	CAGCCCCACC	AGCACCAGCC
6321	CCTCCAGCCC	CACCAGCCCC	AGCCCCTCCT	GTTCCACCGT	GGGTCCCTTT	GCAGCCAATG	CAACTTGGAC	GTTTTTGGGG
6401	TCTCCGGACA	CCATCTCTAT	GTCTTGGCCC	TGATCCTGAG	CCGCCCGGGG	CTCCTGGTCT	TCCGCCTCCT	CGTCCTCGTC
6481	CTCTTCCCCG	TCCTCGTCCA	TGGTTATCAC	CCCCTCTTCT	TTGAGGTCCA	CTGCCGCCGG	AGCCTTCTGG	TCCAGATGTG
6561	TCTCCCTTCT	CTCCTAGGCC	ATTTCCAGGT	CCTGTACCTG	GCCCCTCGTC	AGACATGATT	CACACTAAAA	GAGATCAATA
6641	GACATCTTTA	TTAGACGACG	CTCAGTGAAT	ACAGGGAGTG	CAGACTCCTG	CCCCCTCCAA	CAGCCCCCCC	ACCCTCATCC
6721	CCTTCATGGT	CGCTGTCAGA	CAGATCCAGG	TCTGAAAATT	CCCCATCCTC	CGAACCATCC	TCGTCCTCAT	CACCAATTAC
6801	TCGCAGCCCG	GAAAACTCCC	GCTGAACATC	CTCAAGATTT	GCGTCCTGAG	CCTCAAGCCA	GGCCTCAAAT	TCCTCGTCCC
6881	CCTTTTTGCT	GGACGGTAGG	GATGGGGATT	CTCGGGACCC	CTCCTCTTCC	TCTTCAAGGT	CACCAGACAG	AGATGCTACT
6961	GGGGCAACGG	AAGAAAAGCT	GGGTGCGGCC	TGTGAAGCTA	AGATCTGTCG	ACATCGATGG	GCGCGGGTGT	ACACTCCGCC
7041	CATCCCGCCC	CTAACTCCGC	CCAGTTCCGC	CCATTCTCCG	CCTCATGGCT	GACTAATTTT	TTTTATTTAT	GCAGAGGCCG
7121	AGGCCGCCTC	GGCCTCTGAG	CTATTCCAGA			_		AAAGCTAATT
7001				FI	G 13	(con	td )	

FIG. 13(contd.)

#### **BLASTP v NCBI nr**

Query= INSP037.pep (78 letters)

Database: All non-redundant GenBank CDS

translations+PDB+SwissProt+PIR+PRF

1,446,218 sequences; 465,230,387 total letters

Searching......done

	Score	E
ref   XP_112161.2   similar to putative RNA binding protein 1 [Ratt ref   XP_220945.1   similar to keratin 21, type I, cytoskeletal - r ref   NP_775151.1   cytokeratin 21 [Rattus norvegicus] >gi   125089   s gb   AAD49229.2   AF159462_1 EHEC factor for adherence [Escherichia gb   AAL57562.1   AF453441_46 Efal [Escherichia coli] emb   CAB55629.1   lymphostatin [Escherichia coli] emb   CAC81883.1   Efal-LifA-Tox protein [Escherichia coli] gb   AAA39399.1   ORF1 pir     T36223 hypothetical protein SCE39.13c - Streptomyces coelic >ref   XP_211857.1   hypothetical protein XP_211857 [Homo sapiens]	(bits)	Value
ref XP_211857.1  hypothetical protein XP_211857 [Homo sapiens]	109	8e-24
ref   XP_112161.2   similar to putative RNA binding protein 1 [Ratt	. 38	0.041
ref XP_220945.1  similar to keratin 21, type I, cytoskeletal - r	. 37	0.069
ref NP_775151.1  cytokeratin 21 [Rattus norvegicus] >gi 125089 s	. 37	0.069
gb AAD49229.2 AF159462_1 EHEC factor for adherence [Escherichia	. 35	0.26
gb AAL57562.1 AF453441_46 Efal [Escherichia coli]	35	0.26
emb CAB55629.1  lymphostatin [Escherichia coli]	35	026
emb CAC81883.1  Efal-LifA-Tox protein [Escherichia coli]	35	0.26
gb AAA39399.1  ORF1	35	0.34
pir  T36223 hypothetical protein SCE39.13c - Streptomyces coelic	. 34	0.59
>ref XP_211857.1  hypothetical protein XP_211857 [Homo sapiens]		
Length = 113		

Score = 109 bits (273), Expect = 8e-24Identities = 54/74 (72%), Positives = 63/74 (84%)

Query: 1 MTSPNELNKLPWTNPGETEICDLSDTEFKISVLKNLKEIQDNTEKESRILSDKYKKQIEI 60
MTSPNELN+ P TNP ETEIC++ D EFKI+VL+ L EIQDNTEKE ++LSDK K+IEI

Sbjct: 1 MTSPNELNEAPGTNPAETEICNILDREFKIAVLRKLNEIQDNTEKELKVLSDKIIKEIEI 60

Query: 61 IKGNQAEILELRNA 74

IK NQAEILEL+NA

Sbjct: 61 IKMNQAEILELKNA 74

#### BLAST v month-aa

```
Query= INSP037.pep
         (78 letters)
Database: NCBI: Rolling month (30 days) of new/revised protein
sequences
           37,755 sequences; 14,558,446 total letters
Searching......done
                                                                  Score
                                                                            Ε
                                                                  (bits)
                                                                          Value
Sequences producing significant alignments:
ref | XP 141262.1 | similar to NAG14 protein [Homo sapiens] [Mus mu...
                                                                      30 0.27
ref NP_831679.1 | Phage-related protein [Bacteriophage phBC6A51] ...
                                                                      30 . 0.36
ref|NP 083191.1| RIKEN cDNA 1200008A14 [Mus musculus] >gi|128359...
                                                                      29 0.61
ref NP_852012.1 | neck appendage [Streptococcus phage C1] >gi | 309...
                                                                      28 0.80
ref NP 064648.1 | neurexin I; neurexin I beta; neurexin I alpha; ...
                                                                      28 1.0
                                                                      28 1.0
ref|XP 319358.1| ENSANGP0000006161 [Anopheles gambiae] >gi|2130...
ref|XP 308412.1| ENSANGP0000019827 [Anopheles gambiae] >gi|2129...
                                                                      28 1.0
                                                                      27 1.8
ref NP_196806.2 expressed protein [Arabidopsis thaliana]
gb|AAL29689.1| Snf2-related chromatin remodeling factor SRCAP [T...
                                                                      27 1.8
ref | XP 314825.1 | ENSANGP00000011098 [Anopheles gambiae] >gi | 2129...
                                                                      27 1.8
                                                                      27 2.3
ref|XP 311503.1| ENSANGP00000013657 [Anopheles gambiae] >gi|2129...
>ref | XP_141262.1 | similar to NAG14 protein [Homo sapiens] [Mus musculus]
 ref | XP_230311.1 | similar to NAG14 protein [Homo sapiens] [Rattus norvegicus]
 ref NP 848840.1 RIKEN cDNA 6430556C10 gene [Mus musculus]
 dbj BAC28656.1 unnamed protein product [Mus musculus]
 dbj|BAC33302.1| unnamed protein product [Mus musculus]
          Length = 640
 Score = 30.0 \text{ bits (66)}, Expect = 0.27
 Identities = 22/59 (37%), Positives = 33/59 (55%), Gaps = 8/59 (13%)
Query: 20 ICDLSDTEFK-ISVLKNLKEIQDNTEKESRILSDKYKKQIEIIKGN-----QAEILEL 71
                   K I V KNL+E+ D
                                      +R+L + ++ QI+IIK N
```

Sbjct: 50 VCSCSNQFSKVICVRKNLREVPDGISTNTRLL-NLHENQIQIIKVNSFKHLRHLEILQL 107

#### **FIG. 16A**

#### TBLASTN v NCBI nt-month

```
Ouery= INSP037.pep
         (78 letters)
Database: NCBI: Rolling month (30 days) of new/revised nt sequences
(GenBank+ EMBL + DDBJ sequences (but no EST, STS, GSS, or phase 0, 1
or 2 HTGS sequences))
          44,426 sequences; 216,324,491 total letters
Searching.....done
                                                                           Е
                                                                 Score
                                                                 (bits)
                                                                         Value
Sequences producing significant alignments:
                                                                    105 2e-23
gb|AC093724.3| Homo sapiens BAC clone RP11-1L5 from 2, complete ...
emb|BX510371.4| Human DNA sequence from clone RP13-728A10 on chr...
                                                                     89 2e-18
                                                                     82 4e-16
gb|AC144561.8| Homo sapiens 3 BAC RP11-628C23 (Roswell Park Canc...
dbj|AP001827.5| Homo sapiens genomic DNA, chromosome 11 clone:RP...
                                                                     80 1e-15
emb Z97632.3 HS196E23 Human DNA sequence from clone RP1-196E23 o...
                                                                     66 3e-11
emb|BX322234.7| Human DNA sequence from clone XXyac-65C7 A on ch...
                                                                     62 5e-10
dbj AP005138.3 | Homo sapiens genomic DNA, chromosome 18 clone: RP...
                                                                     54 1e-07
                                                                     54 1e-07
dbi AP006292.2 Homo sapiens genomic DNA, chromosome 9 clone:RP1...
gb|AC083903.10| Homo sapiens chromosome UNK clone RP11-785G23, c...
                                                                     47 1e-05
gb|AY293855.1| Homo sapiens insulin-like growth factor 2 recepto...
                                                                     45 7e-05
>gb|AC093724.3| Homo sapiens BAC clone RP11-1L5 from 2, complete sequence
         Length = 161617
 Score = 105 bits (263), Expect = 2e-23
 Identities = 55/78 (70%), Positives = 62/78 (78%)
 Frame = -3
            MTSPNELNKLPWTNPGETEICDLSDTEFKISVLKNLKEIQDNTEKESRILSDKYKKQIEI 60
Query: 1
            MTSPNELNK P NP ET++CDLS EFKI+VL+ LKEIQDNTEK RILSDK+ K IEI
Sbjct: 22538 MTSPNELNKAPRINPQETKLCDLSHGEFKIAVLRKLKEIQDNTEKGFRILSDKFNKDIEI 22359
            IKGNQAEILELRNADGTL 78
Query: 61
                +AEILEL+NA G L
Sbjct: 22358 IFKTRAEILELKNAIGIL 22305
Score = 30.0 bits (66), Expect = 1.7
 Identities = 19/60 (31%), Positives = 35/60 (57%)
Frame = +3
             NPGETEICDLSDTEFKISVLKNLKEIQDNTEKESRILSDKYKKQIEIIKGNQAEILELRN 73
Ouery: 14
             +P + EI DLS+ EFK+ V+K ++E + E + +
                                                   K + K I + + KG
Sbjct: 111237 DPNKEEITDLSEKEFKL-VIKLIREGPEKGEAQCK----KIQKVIQ*VKGETFKEIDSLN 111401
```

#### TBLASTN v NCBI nt

## **FIG. 16B**

Е

Score

Sequences producing significant alignments: (bits) Value

gb|AC112641.3| Homo sapiens 3 BAC RP11-43118 (Roswell Park Cance... 158 2e-37

gb|AC026118.17| Homo sapiens 3 BAC RP11-67F24 (Roswell Park Canc... 158 2e-37

emb|AL020989.2|HS192P9 Human DNA sequence from clone RP1-192P9 o... 117 3e-25

gb|AC009811.14| Homo sapiens chromosome 3, clone RP11-491K7, com... 116 7e-25

gb|AC108166.5| Homo sapiens BAC clone RP11-724L20 from 4, comple... 115 9e-25

gb|AC011299.3|AC011299 Homo sapiens BAC clone RP11-232C20 from 7... 115 1e-24

gb|AC144613.1| Pan troglodytes chromosome 7 clone RP43-1F6, comp... 115 1e-24

dbj|AP001992.4| Homo sapiens genomic DNA, chromosome 11q clone:R... 115 1e-24

emb|AL359393.9| Human DNA sequence from clone RP11-338I3 on chro... 114 2e-24

emb|AL353577.22| Human DNA sequence from clone RP11-661K19 on ch... 114

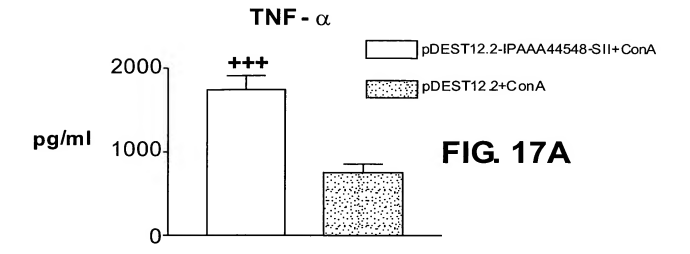
Score = 158 bits (399), Expect = 2e-37
Identities = 78/78 (100%), Positives = 78/78 (100%)
Frame = +3

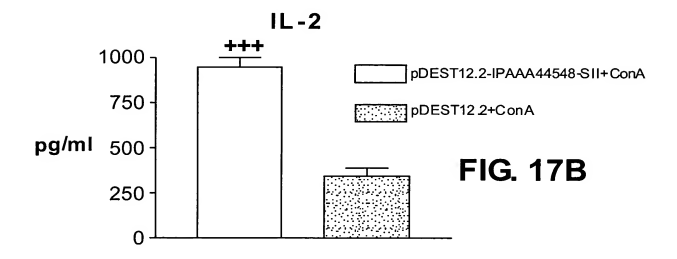
Query: 1 MTSPNELNKLPWTNPGETEICDLSDTEFKISVLKNLKEIQDNTEKESRILSDKYKKQIEI 60 MTSPNELNKLPWTNPGETEICDLSDTEFKISVLKNLKEIQDNTEKESRILSDKYKKQIEI

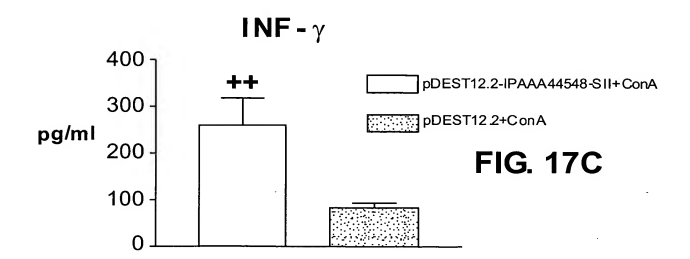
Sbjct: 47052 MTSPNELNKLPWTNPGETEICDLSDTEFKISVLKNLKEIQDNTEKESRILSDKYKKQIEI 47231

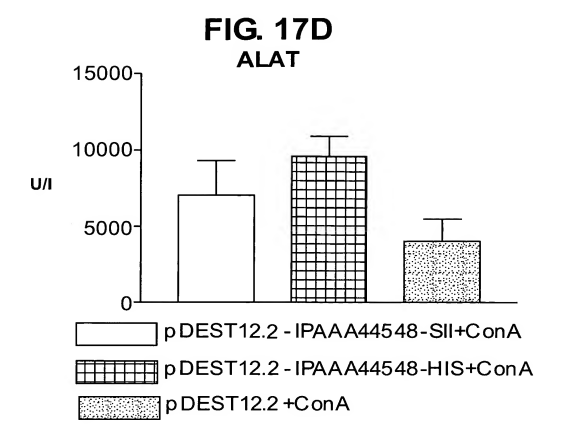
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IKGNQAEILELRNADGTL

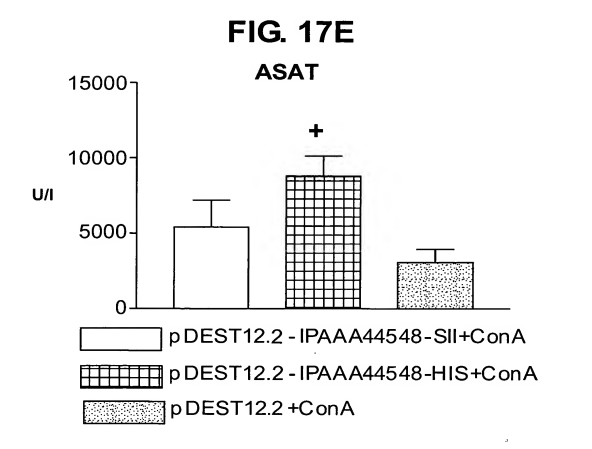
Sbjct: 47232 IKGNQAEILELRNADGTL 47285



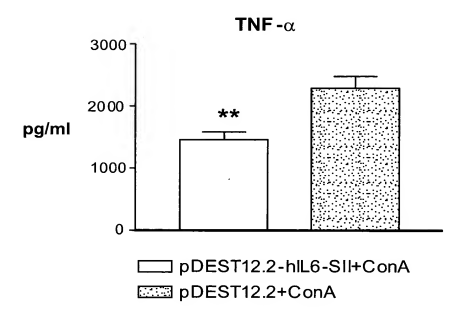




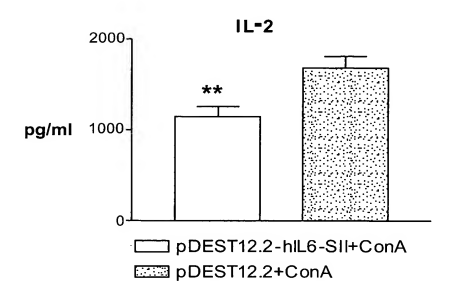




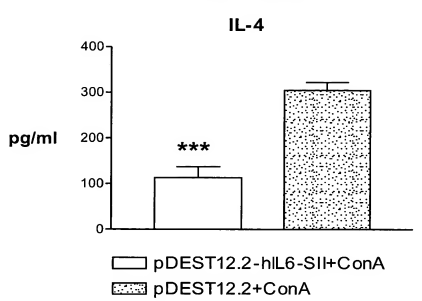
**FIG. 18A** 



**FIG. 18B** 



**FIG. 18C** 



**FIG. 18D** 

